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Usefulness of Genetic Testing in Sudden Cardiac Arrest Survivors With or Without Previous Clinical Evidence of Heart Disease

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Abstract: Genetic testing in survivors of sudden cardiac arrest (SCA) with a suspicious cardiac phenotype is considered clinically useful, whereas its value in the absence of phenotype is disputed. We aimed to evaluate the clinical utility of genetic testing in survivors of SCA with or without cardiac phenotype. Sixty unrelated SCA survivors (median age: 34 [interquartile range 20 to 43] years, 82% male) without coronary artery disease were included: 24 (40%) with detectable cardiac phenotype (Ph(+))SCA after the SCA event and 36 (60%) with no clear cardiac phenotype (Ph(-))SCA. The targeted exome sequencing was performed using the TruSight-One Sequencing Panel (Illumina). Variants in 185 clinically relevant cardiac genes with minor allele frequency <1% were analyzed. A total of 32 pathogenic or likely pathogenic variants were found in 27 (45%) patients: 17 (71%) in the Ph(+))SCA group and 10 (28%) in the Ph(-))SCA group. Sixteen (67%) Ph(+))SCA patients hosted mutations congruent with the suspected phenotype, in which 12 (50%) were cardiomyopathies and 4 (17%) channelopathies. In Ph(-))SCA cases, 6 (17%) carried a mutation in cardiac ion channel genes that could explain the event. The additional 4 (11%) mutations in this group, could not explain the phenotype and require additional studies. In conclusion, cardiac genetic testing was positive in nearly 2/3 patients of the Ph(+))SCA group and in 1/6 of the Ph(-))SCA group. The test was useful in both groups to identify or confirm an inherited heart disease, with an important impact on the patient care and first-degree relatives at risk.

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Usefulness of Genetic Testing in Sudden Cardiac Arrest Survivors With or Without Previous Clinical Evidence of Heart Disease

Short title: Genetic Basis of Sudden Cardiac Arrest

Babken Asatryan, MD^a, André Schaller, PhD^b, Jens Seiler, MD^a, Helge Servatius, MD^a, Fabian Noti, MD^a, Samuel H. Baldinger, MD^a, Hildegard Tanner, MD^a, Laurent Roten, MD^a, Roger Dillier, MD^c, Anna Lam, MD^{a,d}, Andreas Haeberlin, MD, PhD^a, Giulio Conte, MD, PhD^e, Ardan M. Saguner, MD^f, Stephan Andreas Müller, MD^c, Firat Duru, MD^f, Angelo Auricchio, MD, PhD^e, Peter Ammann, MD^g, Christian Sticherling, MD^h, Haran Burri, MDⁱ, Tobias Reichlin, MD^a, Matthias Wilhelm, MD^a, Argelia Medeiros-Domingo, MD, PhD^{a,*}

^a Department of Cardiology, Inselspital, Bern University Hospital, University of Bern, Switzerland

^b Division of Human Genetics, Department of Pediatrics, Inselspital, Bern University Hospital, University of Bern, Switzerland

^c Stadtspital Triemli, Zurich, Switzerland

^d Hôpital Cardiologique du Haut-Lévêque - CHU de Bordeaux, Bordeaux-Pessac, France

^e Division of Cardiology, Fondazione Cardiocentro Ticino, Lugano, Switzerland

^f University Heart Center Zurich, Zurich, Switzerland

^g Kantonsspital, St. Gallen, Switzerland

^h Department of Cardiology, University Hospital Basel, Basel, Switzerland

ⁱ University Hospital of Geneva, Geneva, Switzerland

*Corresponding author:

Argelia Medeiros-Domingo

Department of Cardiology, Inselspital, 3010 Bern, Switzerland.

Phone: +41 31 632 03 63 Fax: +41 31 632 14 14

E-mail: argelia.medeiros@hin.ch

Declaration of interest

Dr. Saguner reports personal fees from Boston Scientific, and grants from Biosense Webster, Biotronik, Abbott, and BMS Pfizer, outside the submitted work. Dr. Müller reports personal fees from Biotronik, Medtronic, St. Jude Medical/ Abbott, Biosense Webster, AstraZeneca, Daiichi Sankyo, LivaNova/ Sorin, and MSD, and training support from Biotronik, Boston Scientific, Medtronic, and Biosense Webster, outside the submitted work. Prof. Auricchio serves as a consultant to Abbott, Biosense Webster, Daiichi-Sankyo, Boston Scientific, Cardiotek-Schwarzer, Cordis Biologics Delivery Systems, and Medtronic. He also received speaker fees from Abbott, Biosense-Webster, Daiichi-Sankyo, Boston Scientific, and Medtronic. Other authors have no conflicts of interests to declare.

Abstract

Genetic testing in survivors of sudden cardiac arrest (SCA) with a suspicious cardiac phenotype is considered clinically useful, whereas its value in the absence of phenotype is disputed. We aimed to evaluate the clinical utility of genetic testing in survivors of SCA with or without cardiac phenotype. Sixty unrelated SCA survivors (median age: 34 [IQR 20-43] years, 82% male) without coronary artery disease were included: 24 (40%) with detectable cardiac phenotype (Ph(+))SCA) after the SCA event and 36 (60%) with no clear cardiac phenotype (Ph(-))SCA). Either targeted exome sequencing was performed using the TruSight-One Sequencing Panel (Illumina). Variants in 185 clinically relevant cardiac genes with minor allele frequency <1% were analyzed. A total of 32 pathogenic or likely pathogenic variants were found in 27 (45%) patients: 17 (71%) in the Ph(+))SCA group and 10 (28%) in the Ph(-))SCA group. Sixteen (67%) Ph(+))SCA patients hosted mutations congruent with the suspected phenotype, in which 12 (50%) were cardiomyopathies and 4 (17%) channelopathies. Among Ph(-))SCA cases, 6 (17%) carried a mutation in cardiac ion channel genes that could explain the event. The additional 4 (11%) mutations in this group, could not explain the phenotype and require additional studies. In conclusion, cardiac genetic testing was positive in nearly 2/3 patients of the Ph(+))SCA group and in 1/6 of the Ph(-))SCA group. The test was useful in both groups to identify or confirm an inherited heart disease, with an important impact on the patient care and first-degree relatives at risk.

Keywords: idiopathic ventricular fibrillation; cardiac channelopathy; cardiomyopathy; genetic testing; sudden cardiac death

INTRODUCTION

Sudden cardiac arrest (SCA) is a common manifestation of a wide spectrum of inherited and acquired cardiovascular diseases, and extra-cardiac pathologies.¹ Identifying the cause of SCA is essential for proper management of the arrhythmic substrate to allow a personalized approach. The detection of a specific cardiomyopathy or inherited arrhythmia phenotype in a SCA survivor (Ph(+)-SCA) allows genetic testing for confirmation of the clinical diagnosis, risk stratification, and eventually specific therapy.^{2, 3} Phenotype-guided genetic testing has shown a heterogeneous diagnostic yield of 9-60% in SCA survivors, depending on the panel and technology used.⁴⁻⁶ In the absence of a clear cardiac phenotype, the event is referred to as a phenotype-negative SCA (Ph(-)-SCA), or idiopathic ventricular fibrillation.⁷ Genetic testing in Ph(-)-SCA has been discouraged by the expert consensus (class III) to avoid possible over-diagnosis and unnecessary interventions in families owing to potential misinterpretation of innocent bystander variants;^{2, 7} Thus, currently the knowledge on genetic basis of Ph(-)-SCA is limited. In this study, we sought to evaluate the clinical and genetic characteristics of an unselected cohort of SCA survivors referred for genetic testing from different regions of Switzerland.

METHODS

In this partly prospective and partly retrospective study we included SCA patients referred to the Bern University Hospital for genetic testing between January 2014 and May 2018. SCA was defined as an unexpected circulatory arrest with cardiovascular collapse and documented hemodynamically unstable sustained ventricular tachycardia (VT) or ventricular fibrillation (VF), restored by cardioversion or defibrillation, as appropriate. Internationally accepted criteria were used to diagnose/exclude specific diseases. Initial diagnostic workup included personal and family history, 12-lead resting ECG, echocardiography and coronary angiogram/ cardiac

computed tomography, exercise stress test (whenever possible); additional examination with cardiac MRI, electrophysiological study, or provocative testing were performed when considered clinically indicated. Patients with coronary artery disease, myocarditis, sarcoidosis, systemic diseases affecting the heart, secondary cardiomyopathies, or reversible causes of SCA were excluded from the study. Survivors of SCA with any suspicious or robust cardiac phenotype (Ph(+)-SCA) and those with no identified phenotype (Ph(-)-SCA) were included in this study. This study was conducted in full agreement with the principles outlined in the Declaration of Helsinki. The study protocol was approved by the Ethical Commission of the Canton Bern.

Genomic DNA was extracted from EDTA-blood using standard procedures. Clinical exome analysis was performed using next-generation sequencing on MiSeq or NextSeq 500 instrument using either the TruSight One or TruSight One expanded Sequencing Panel (Illumina, San Diego, CA), which provide comprehensive coverage of exons and flanking intronic sequences of 4813 and 6794 clinically relevant genes, respectively. The intronic coverage comprised at least 30 to 50bp of each intron; the coverage depth of exons was minimum 30x. After sequencing, variants in 185 genes associated with hereditary cardiovascular diseases were analyzed (Supplementary Table 1). Read alignment and local realignment of indels were performed using CLC Genomic Workbench v7.5.1 or CLC Biomedical Genomic Workbench v5.0 (Qiagen, Redwood City, CA). Novel or putative disease-associated variants were analyzed using the following filtering criteria: protein's primary structure, species conservation of the affected amino acid and an allele frequency <1% based on the 1000 Genome Project database. Exonic variants below 1% frequency in the Exome Aggregation Consortium browser, including +/- 8bp flanking intronic variants, were further analyzed. For titin (*TTN*), only radical mutations (i.e. nonsense, frameshift, and splice-site mutations) were considered as putative pathogenic variants. The final interpretation of variants was performed according to the guidelines

established by the American College of Medical Genetics and Genomics (ACMG).⁸

Categorical variables are presented as numbers and percentages; quantitative variables are presented as median (interquartile range [IQR]) for non-normally distributed variables. Categorical variables were compared using Chi-Square or Fisher's exact test, as appropriate. Continuous variables were compared using Student's *t* test or Mann-Whitney *U* test, as appropriate. To adjust for multiple testing, we used Bonferroni adjustment and set alpha at 0.005. Statistical analyses were performed using IBM SPSS Statistics for Windows, Version 25.0. IBM Corp. All tests were 2-tailed, with a $p < 0.05$ considered significant.

RESULTS

Sixty unrelated SCA patients (median age 34 [20-43] years; male-82%) without previously known heart disease, including 24 (40%) with (Ph+)SCA and 36 (60%) with Ph(-)SCA, were enrolled (Table 1). The two groups were similar in terms of age at SCA and gender distribution. Data regarding the circumstances of SCA were available in 28 (47%) patients. In 13 (22%) cases the event occurred at the time or shortly after exercise activities and in 15 (25%) occurred unrelated to sports activities. The proportion of exercise-related SCA cases were similar in both groups (19% vs 25%, $p = 0.609$). Twelve (92%) out of 13 exercise-related SCA cases occurred in males, but males were not more likely to experience a SCA during exercise, than the females ($p = 0.263$).

Overall, 32 putative pathogenic variants (14 novel and 18 previously reported) were identified in 27 (45%) out of 60 subjects (Table 2). No variants have been identified more than once in our cohort. Patients with exercise-related SCA had a likelihood of having a positive result of genetic testing (54%) similar to those with exercise-unrelated (40%) or unknown circumstances of SCA (44%; $p = 0.747$).

In the Ph(+)SCA group, dilated cardiomyopathy (DCM, $n = 7$) was the most common,

followed by hypertrophic cardiomyopathy (n=4) and arrhythmogenic right-ventricular cardiomyopathy (ARVC, n=4). Other phenotypes included left-ventricular non-compaction, catecholaminergic polymorphic ventricular tachycardia (CPVT, n=2), mitral valve prolapse (n=2), long QT syndrome (n=1), short QT syndrome (n=1), and progressive cardiac conduction disease (n=1). Twenty-one putative pathogenic variants were identified in 17 (71%) out of 24 Ph(+)-SCA patients, including 14 (67%) mutations in cardiomyopathy-associated genes (*LMNA*, *TNNT2*, *TNNI3*, *TTN*, *FLNC*, *PKP2*, *DSP*, *JPH2*, *TNXB*, *LAMP2*, *BAG3*, and *CAV3*) and 7 (33%) in cardiac ion channel genes (*RYR2*, *CASQ2*, *SCN5A*, *KCNQ1*, and *HCN4*). In 13 out of 17 (76%) genotype-positive patients a cardiomyopathy and in 4 (24%) cardiac channelopathy was suspected (Figure 1). Sixteen (67%) out of 24 Ph(+)-SCA patients carried mutations in agreement with the observed phenotype, whereas 1 patient, a 31-year-old male, carried a likely pathogenic variant in the *CAV3* (caveolin 3) gene, which so far has not been associated with the identified DCM phenotype; thus it was considered suspicious but not diagnostic.

Overall, 11 pathogenic or likely pathogenic variants were detected in 10 (28%) out of 36 Ph(-)-SCA survivors (Figure 1), which was significantly lower than the 71% diagnostic yield in Ph(+)-SCA patients ($p=0.017$). Six (17%) subjects carried mutations in ion channel genes (*SCN5A*, *RYR2*, and *KCNQ1*), that could explain the SCA in the absence of phenotype (concealed channelopathies). Additionally, 5 putative pathogenic variants localized in the *RANGRF*, *ABCC9*, *TRPM4*, *DSP* and *TTN* did not clarify the SCA event and require further functional studies or segregation analysis.

DISCUSSION

In a cohort of 60 SCA survivors without previously known heart disease, the cardiac panel genetic testing detected 32 putative pathogenic variants in 27 (45%) cases. In patients with an identifiable cardiac phenotype, the genetic test could explain the phenotype and the SCA in 16

(67%) cases, while in those with no phenotype, the genetic test could clarify the cause of the SCA in 6 (17%) cases. Approximately 20% of all patients with SCA were found to have an association with sports activities or physical work.

Our Ph(+)SCA cohort included cases with a wide variety of cardiac phenotypes, which once again highlights the broad spectrum of heritable heart diseases underlying SCA. As expected, the diagnostic yield was 2-fold higher in the Ph(+)SCA group than in the Ph(-)SCA. Interestingly, the diagnostic yield of 71% in the Ph(+)SCA group surpassed the reported general yield for each of the diseases suspected in these patients probably because stronger phenotypes are usually associated with higher diagnostic yield of the genetic test. All but one of the genotype-positive cases of the Ph(+)SCA group, carried pathogenic variants consistent with the suspected heart disease, underscoring that phenotyping remains central to the diagnosis of the cause of SCA. Clarifying the clinical role of the potentially disease-causing variants in genes unrelated to the phenotypes, such as the *CAV3*-variant in a patient with DCM, requires further research and highlights the utility of broader genetic panels for research purposes.

The possible genetic nature of SCA in the absence of measurable heart disease has been first recognized in 2000, when the p.Ser1710Leu mutant sodium channel has been shown to be the likely molecular substrate for Ph(-)SCA.⁹ Subsequently, detailed experimental and familial studies have established the causality for the Dutch founder haplotype in *DPP6*,¹⁰ mutations in *CALM1*,¹¹ *IRX3*,¹² and *RYR2*.¹³ Moreover, recent studies on Ph(-)SCA cohorts found mutations in cardiac ion channel genes (*RYR2*, *KCNH2*, *KCNQ1*, *KCNE1*, *SCN5A*, and *CACNA1C*) and cardiomyopathy-associated genes (*LMNA*, *TTN*, *DSP*, *PLN*, *DSC2*, *MYBPC3*, *MYH7*, *TNNT2*), with a diagnostic yield of up to 15%, thus further expanding the spectrum of genes potentially associated with Ph(-)SCA.^{3, 14} We identified a genetic substrate that explains nearly one-sixth of our Ph(-)SCA cases. More than half of these mutations were located in principal cardiac ion

channel genes. This yield is consistent with previous reports,^{3, 14, 15} and highlights that concealed cardiac channelopathies may manifest with malignant arrhythmias even in the absence of a detectable cardiac phenotype.

Four out of 6 mutations detected in our Ph(-)SCA cohort were located in *RYR2*, making CPVT the most prevalent diagnosis in this group (11%). Two CPVT patients (#3 and #9) had a history of syncopal episodes preceding SCA with no direct link to exercise. Another patient (#5) had frequent premature ventricular contractions and a non-sustained VT at maximum exercise load, which did not raise a suspicion of CPVT. The fourth patient (#6) had a negative exercise test but experienced a VT at follow up terminated with an ICD shock. These data, in line with recent reports,¹⁵ indicate the need for a low threshold to suspect CPVT and to perform a confirmatory genetic test in Ph(-)SCA survivors with equivocal stress test findings.

The remaining putative pathogenic variants affected genes encoding the ion channel TRPM4 (*TRPM4*), ion channel interacting proteins (*RANGRF*, *ABCC9*) or myocardial structural proteins (*DSP*, *TTN*). Mutations in the TRPM4 channel gene have been implicated in progressive familial heart block, atrioventricular block, right bundle-branch block, and Brugada syndrome (BrS).¹⁶ Although the TRPM4 channel is known to be involved in cellular calcium regulation, its exact role in the cardiac physiology and pathophysiology remains unclear. Loss-of-function *RANGRF* mutations have been reported to cause BrS through impairing the trafficking of Nav1.5 channel.¹⁷ Likewise, gain-of-function mutations in *ABCC9*, a critical gene for the activation and regulation of the K_{ATP} channels, were shown to result in increased I_{KATP} current and manifest with BrS.¹⁸ Mutations in *DSP* (desmoplakin) have been implicated in ARVC¹⁹ and DCM.²⁰ Desmoplakin disease has been shown to cause connexin mislocalization and result in significant alterations in conduction-repolarization abnormalities prior to morphological changes.²¹ Radical mutations in *TTN* have been linked mainly to DCM,²³ with overlaps with other cardiomyopathies

and are known to confer high arrhythmic potential. Despite the presumed pathogenic role of these variants, their potential causality in Ph(-)SCA is subject to further evaluation in functional and follow-up studies. These patients would therefore need close surveillance for detection of any related phenotype at follow-up.

Our findings are particularly interesting in the context of the current HRS/ EHRA guidelines, which discourage the use of genetic testing in Ph(-)SCA.^{2, 7} On the contrary, consideration of genetic testing in SADS is recommended (IIa).^{2, 7} The main difference between these two groups is that survivors reach the hospital alive and SADS victims to the morgue. Previous studies have found a genetic cause in 25% of SADS cases.^{24, 25} In such cases, cascade screening reveals a cardiac disease in nearly half of relatives,⁶ who can benefit from regular surveillance, lifestyle recommendations, and preventive therapy. Our results indicate that the yield of the genetic test in survivors of Ph(-)SCA (17%) is similar to those in SADS victims. Additionally, our analysis uncovered cardiomyopathy gene mutations in Ph(-)SCA patients, which is in accordance with studies on SADS and requires further studies.²⁵ Predictive genetic testing in Ph(-)SCA patients' family members can therefore be as useful as in relatives of SADS victims.^{2, 7}

Similar to previous studies, the use of large gene panels in our study led to high yield of VUS, particularly in the Ph(-)SCA group (47%). The clinical implication of these variants remain elusive and require further studies. Mutations in minor cardiomyopathy or channelopathy genes, defined as implicated in <5% of disease cases, were identified in 8 (13%) SCA patients. These included mutations in *RANGRF*, *ABCC9*, *FKTN*, *KCNE2* in patients with Ph(-)SCA and *LAMP2*, *BAG3*, *HCN4*, *CAV3*, *FLNC*, and *JPH2* in patients with Ph(+)SCA.

Our results suggest that genetic testing is useful in all SCA survivors with and without evidence of cardiac disease. Nevertheless, the test should be performed only by experienced and

accredited laboratories to avoid potential erroneous assignment of pathogenicity.³ Longitudinal prognostic studies of a large cohort of genetically tested SCA survivors and their family members are necessary to determine the potential role of mutation status for risk stratification. The progress in the understanding of the genotype-phenotype associations and the evolving variant classification schemes will optimistically solve the puzzle of uncertain variants and provide more reliable criteria for assessment of pathogenicity

Our study has several limitations. The number of enrolled patients in the present study was relatively small, an inherent problem in studies conducted on rare diseases. Our cohort included only SCA survivors referred for genetic testing. Although the classification of the variants stringently followed the ACMG guidelines, we cannot definitely conclude on their pathogenicity in the absence of segregation or functional studies, although this may be hampered by an incomplete penetrance of the variants. We did not investigate whether a carrier status of a putative pathogenic variant confers a risk for recurrent cardiac events, or whether mutation-carrier Ph(-)SCA patients develop a phenotype at follow-up. Finally, we lack clinical and genetic data of the families, which could help better interpret the impact of many variants.

In conclusion, comprehensive cardiac panel genetic testing identified mutations linked to channelopathies and/or cardiomyopathies in nearly half of SCA survivors without previous diagnosis of cardiac disease. Genetic testing revealed a putative pathogenic variant compatible with the phenotype in 67% of Ph(+)SCA cases. In the Ph(-)SCA group, genetic test revealed the cause of arrest in 17% of cases, similar to the yield reported in SADS. Majority of mutations identified in Ph(-)SCA patients were associated with channelopathies, most commonly CPVT, whereas in the Ph(+)SCA group, the mutations were mostly located in cardiomyopathy-associated genes. Since genetic testing facilitated the understanding of one-sixth of Ph(-)SCA cases, the test is recommended in all SCA survivors, with or without a cardiac phenotype, as it

allows to guide treatment decisions. This study also highlights the need for long-term follow-up studies to examine the prognostic value of genetic findings in SCA survivors.

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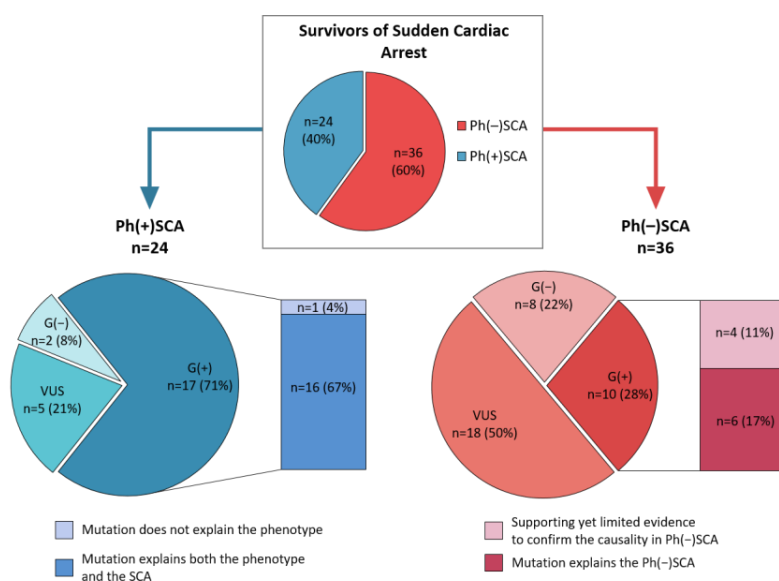


Figure 1. The results and the interpretation of genetic test findings in survivors of sudden cardiac arrest with and without evident clinical cardiac phenotype. G, genotype; Ph, phenotype; SCA, sudden cardiac arrest; VUS, variant of uncertain significance.

Table 1. Demographic characteristics and genetic test results in sudden cardiac arrest survivors referred for cardiovascular genetic testing, grouped based on the displayed cardiac phenotype.

Characteristic	All patients with SCA	Patients with Ph(+)-SCA	Patients with Ph(-)-SCA	<i>P</i> -value *
No. of probands	60 (100%)	24 (40%)	36 (60%)	0.155
Age at SCA, y, median (IQR)	34 (20-43)	36 (20-46)	34 (20-42)	0.490
Male gender	49 (82%)	21 (87%)	28 (78%)	0.274
Patients aged ≤ 35 yrs	32 (53%)	12 (50%)	20 (56%)	0.673
Male	27 (84%)	10 (83%)	17 (85%)	0.674
Female	5 (16%)	2 (17%)	3 (15%)	
Genetic test results				
Positive	27 (45%)	17 (71%)	10 (28%)	0.001

Male	23 (85%)	15 (88%)	8 (80%)	0.561
Female	4 (15%)	2 (12%)	2 (20%)	
VUS	23 (38%)	5 (21%)	18 (50%)	0.709
Male	18 (78%)	4 (80%)	14 (78%)	
Female	5 (22%)	1 (20%)	4 (22%)	0.622
Negative	10 (17%)	2 (8%)	8 (22%)	
Male	8 (80%)	2 (100%)	6 (75%)	0.622
Female	2 (20%)	0 (0%)	2 (25%)	

Abbreviations: Ph, phenotype; SCA, sudden cardiac arrest; SD, standard deviation; VUS, variant of uncertain significance.

* P values are for binomial test, t-test, Fisher's exact test, or Chi-Square test, as appropriate. Chi-Square test was used to test for gender differences between phenotype groups in the whole population, as well as in subgroups according to age and genetic test results.

Table 2. Summary of clinical and genetic characteristics of sudden cardiac arrest survivors.

Case #	Gender	Age at SCA, y	Phenotype category	Cardiac Phenotype	Circumstances of the event	Gene	Reference sequence no.	Nucleotide change	Protein change	Variant type	Homo- Heterozygous	dbSNP ID	AF in gnomAD (European non-Finnish)	MAF in ExAC	ACMG variant class
1	M	56	Ph(-)-SCA	-	exercise unrelated	<i>RANGRF</i>	NM_16492.4	c.198C>A	p.Tyr68*	nonsense	heterozygous	rs1223777741	0.00001758	N/A	LP
2	M	17	Ph(-)-SCA	-	unknown	<i>DES</i>	NM_001927.3	c.742C>T	p.Arg248Cys	missense	heterozygous	rs772117708	0.000008920	0.000008	VUS
						<i>SCN5A</i>	NM_001099404.1	c.2441G>A	p.Arg814Gln	missense	heterozygous	rs199473584	0.00003158	0.000018	LP
3	F	5	Ph(-)-SCA	-	exercise unrelated	<i>RYR2</i>	NM_001035.2	c.6737C>T	p.Ser189Leu	missense	heterozygous	rs144860423	0.0003642	0.0002	VUS
4	M	17	Ph(-)-SCA	-	exercise-related	<i>ABCC9</i>	NM_020297.3	c.1320+1G>A	p.Ser2246Leu	missense	heterozygous	rs121918597	N/A	N/A	P
5	F	17	Ph(-)-SCA	-	exercise unrelated	<i>RYR2</i>	NM_001035.2	c.365G>A	p.(?)	splice region	heterozygous	rs139620148	0.00003899	0.000058	LP
6	M	47	Ph(-)-SCA	-	unknown	<i>RYR2</i>	NM_001035.2	c.1250G>A	p.Arg122His	missense	heterozygous	rs727503396	N/A	N/A	LP
7	M	28	Ph(-)-SCA	-	exercise unrelated	<i>DSP</i>	NM_004415.2	c.4775A>G	p.Arg417Gln	missense	heterozygous	rs751428303	0.00001559	0.000025	LP
8	F	34	Ph(-)-SCA	-	unknown	<i>TTN</i>	NM_001256850	c.29732-2A>T	p.Lys1592Arg	missense	heterozygous	rs200421954	0.0003817	0.000185	LP
						<i>TRPM4</i>	NM_017636	c.238T>C	p.?	splice region	heterozygous	novel variant	N/A	N/A	P
9	M	38	Ph(-)-SCA	-	exercise-related	<i>FKTN</i>	NM_001079802	c.1380dupA	p.Leu796Pro	missense	heterozygous	rs1191927360	N/A	N/A	VUS
						<i>RYR2</i>	NM_001035.2	c.1346T>G	p.Tyr461Ilefs*?	frameshift	heterozygous	rs775366895	0.00004399	N/A	LP*
10**	M	2	Ph(-)-SCA	-	unknown	<i>DSP</i>	NM_004415.2	c.2462A>G	p.Ile449Arg	missense	heterozygous	rs373331669	0.00001777	0.000008	LP
11	M	28	Ph(-)-SCA	-	unknown	<i>KCNE2</i>	NM_172201.1	c.170T>C	p.Lys821Arg	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>CACNB2</i>	NM_201593.3	c.410G>A	p.Ile57Thr	missense	heterozygous	rs74315448	0.001045	0.0009	P†
12**	M	22	Ph(-)-SCA	-	exercise unrelated	<i>SCN5A</i>	NM_001099404.1	c.3929C>T	p.Gly137Asp	missense	heterozygous	rs754596850	0.000008799	0.000008	VUS
						<i>RYR2</i>	NM_001035.2	c.7570G>A	p.Pro1310Leu	missense	heterozygous	novel variant	N/A	N/A	VUS
13	M	19	Ph(-)-SCA	-	exercise unrelated	<i>TTN</i>	NM_001267550.1	c.1545_1549delAGAAA	p.Val2524Ile	missense	heterozygous	rs934248102	0.00003893	N/A	VUS
						<i>KCNQ1</i>	NM_000218.2	c.190_210delCCTGCGTCCCCGGCGCGCGCC	p.Lys515AsnfsTer2	frameshift	heterozygous	novel variant	N/A	N/A	LP
14	F	41	Ph(-)-SCA	-	exercise-related	<i>PKP2</i>	NM_004572.3	c.2431C>A	p.Pro64_Pro70del	deletion	heterozygous	reported, no dbSNP ID	N/A	N/A	VUS‡
						<i>AKAP9</i>	NM_005751	c.1066A>T	p.Arg811Ser	missense	heterozygous	rs139734328	0.0009214	0.00087	VUS
15	M	28	Ph(-)-SCA	-	unknown	<i>TNXB</i>	NM_019105.6	c.5956A>G	p.Asp3555Val	missense	heterozygous	rs139046510	0.001444	0.0009	VUS
						<i>TNNI3</i>	NM_000363	c.349A>G	p.Ile1986Val	missense	heterozygous	rs201365475	0.000008873	N/A	VUS
16	M	24	Ph(-)-SCA	-	unknown	<i>DSP</i>	NM_004415.2	c.2723G>A	p.Lys117Glu	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>MYH6</i>	NM_002471.3	c.1138G>A	p.Arg908His	missense	heterozygous	rs142494121	0.0009762	0.0011	VUS
17	M	35	Ph(-)-SCA	-	exercise-related	<i>CTNNA3</i>	NM_001127384	c.2573T>C	p.Glu380Lys	missense	heterozygous	rs768924353	N/A	0.000008	VUS
						<i>KCNE1</i>	NM_001270402.1	c.253G>A	p.Leu858Ser	missense	heterozygous	rs41313840	0.0003953	0.0003	VUS
18	M	56	Ph(-)-SCA	-	exercise-related	<i>KCNH2</i>	NM_000238	c.1653C>G	p.Asp85Asn	missense	heterozygous	rs1805128	0.01223	0.0092	VUS
						<i>FLNC</i>	NM_001458.4	c.1616C>T	p.Phe551Leu	missense	heterozygous	novel variant	N/A	N/A	VUS
19	M	41	Ph(-)-SCA	-	unknown	<i>ANK2</i>	NM_001148.4	c.4799T>C	p.Pro539Leu	missense	heterozygous	rs375570393	0.000	0.000017	VUS
						<i>SCN2B</i>	NM_004588.4	c.356C>T	p.Ile1600Thr	missense	heterozygous	rs764150279	0.00002342	0.000033	VUS
20	M	38	Ph(-)-SCA	-	unknown	<i>KCNJ2</i>	NM_000891.2	c.973C>T	p.Pro119Leu	missense	heterozygous	rs767589740	N/A	0.000200	VUS
21	M	31	Ph(-)-SCA	-	unknown	<i>KCNJ5</i>	NM_000890	c.155G>A	p.Arg325Cys	missense	heterozygous	rs202067116	0.00006197	0.000041	VUS
22	M	34	Ph(-)-SCA	-	exercise unrelated	<i>FBN1</i>	NM_000138	c.1307A>G	P.Arg52His	missense	heterozygous	rs144062083	0.00004396	0.000025	VUS
						<i>SCN10A</i>	NM_006514.3	c.3675T>G	p.Tyr436Cys	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>DSG2</i>	NM_001943.3	c.1642C>T	p.Ile1225Met	missense	heterozygous	rs371834340	0.00008519	0.000017	VUS
									p.Arg548Cys	missense	heterozygous	rs550400909	0.00003125	0.000009	VUS

23	M	46	Ph(-)-SCA	-	exercise-related	<i>KCNE1</i>	NM_000219.4	c.314C>T	p.Ser105Leu	missense	heterozygous	rs780041404	0.00002639	0.000091	VUS
						<i>RYR2</i>	NM_001035.2	c.11018G>A	p.Arg3673Gln	missense	heterozygous	rs778111237	0.00003552	0.000025	VUS
						<i>SCN1B</i>	NM_199037	c.716C>A	p.Thr239Asn	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>IDH2</i>	NM_002168.3	c.293T>C	p.Ile98Thr	missense	heterozygous	rs139512088	0.00007035	0.000033	VUS
24	F	57	Ph(-)-SCA	-	exercise unrelated	<i>MYH7</i>	NM_000257.3	c.3078G>T	p.Lys1026Asn	missense	heterozygous	novel variant	N/A	N/A	VUS
25	M	48	Ph(-)-SCA	-	unknown	<i>MYH6</i>	NM_002471.3	c.4429C>T	p.Arg1477Cys	missense	heterozygous	rs201989347	0.00006166	0.000017	VUS
26	M	33	Ph(-)-SCA	-	unknown	<i>KCNH2</i>	NM_000238.3	c.3304A>C	p.Thr1102Pro	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>KCNH2</i>	NM_000238.3	c.3265A>C	p.Thr1089Pro	missense	heterozygous	novel variant	N/A	N/A	VUS
27	M	24	Ph(-)-SCA	-	exercise unrelated	<i>TMEM43</i>	NM_024334.2	c.323T>C	p.Val108Ala	missense	heterozygous	rs182351748	0.000	0.0002	VUS
						<i>MYH6</i>	NM_002471.3	c.982G>A	p.Glu328Lys	missense	heterozygous	rs144422878	0.00007752	0.000058	VUS
28	F	39	Ph(-)-SCA	-	unknown	<i>SCN10A</i>	NM_006514	c.1355C>T	p.Ser452Phe	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>SNTA1</i>	NM_003098	c.692C>T	p.Pro231Leu	missense	heterozygous	rs559396761	0.000	0.000008	VUS
						<i>TNXB</i>	NM_019105	c.12373G>A	p.Glu4125Lys	missense	heterozygous	novel variant	N/A	N/A	VUS
29	M	60	Ph(-)-SCA	-	unknown	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
30	M	13	Ph(-)-SCA	-	unknown	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
31	F	39	Ph(-)-SCA	-	unknown	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
32	M	19	Ph(-)-SCA	-	exercise-related	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
33	M	18	Ph(-)-SCA	-	unknown	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
34	M	36	Ph(-)-SCA	-	exercise unrelated	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
35	M	42	Ph(-)-SCA	-	unknown	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
36**	F	43	Ph(-)-SCA	-	unknown	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
37	F	37	Ph(+)-SCA	borderline ARVC	unknown	<i>RYR2</i>	NM_001035.2	c.12665_12667delAGA	p.Lys4223del	deletion	heterozygous	novel variant	N/A	0.000091	LP
38	F	24	Ph(+)-SCA	LVNC/ DCM	exercise unrelated	<i>LAMP2</i>	NM_002294.2	c.928+1G>T	p.(?)	splice region	heterozygous	novel variant	N/A	N/A	P
39	M	62	Ph(+)-SCA	LQTS	exercise unrelated	<i>KCNQ1</i>	NM_000218.2	c.691C>T	p.Arg231Cys	missense	heterozygous	rs199473457	0.00006497	N/A	LP
40	M	15	Ph(+)-SCA	suspected CPVT	unknown	<i>RYR2</i>	NM_001035.2	c.6741C>T	p.Thr2914Met	missense	heterozygous	rs766484260	0.00001773	0.000035	VUS
								c.12257G>A	p.Arg4086His	missense	heterozygous	rs752977469	0.000008947	0.000009	LP
41	M	19	Ph(+)-SCA	DCM/ Laminopathy	unknown	<i>BAG3</i>	NM_004281	c.803C>A	p.Ser268*	nonsense	heterozygous	novel variant	N/A	N/A	P
42	M	69	Ph(+)-SCA	possible HCM	unknown	<i>TNNI2</i>	NM_001001430.2	c.860G>A	p.Trp287*	nonsense	heterozygous	rs727504247	0.000	0.000011	P
43	M	41	Ph(+)-SCA	PCCD	unknown	<i>SCN5A</i>	NM_198056.2	c.3840+1G>A	p.(?)	splice region	heterozygous	rs1366120635	0.000008801	N/A	P
						<i>C7NMA3</i>	NM_013266.2	c.2097C>G	p.Asn699Lys	missense	heterozygous	rs142942346	N/A	N/A	VUS
						<i>ANK2</i>	NM_001148.5	c.10825C>A	p.Leu3609Ile	missense	heterozygous	novel variant	N/A	N/A	VUS
44	M	45	Ph(+)-SCA	LVNC	unknown	<i>HCN4</i>	NM_005477.2	c.1435A>G	p.Ile479Val	missense	heterozygous	novel variant	N/A	N/A	P
								c.1454C>A	p.Ala485Glu	missense	heterozygous	novel variant	N/A	N/A	P
45	M	22	Ph(+)-SCA	CPVT	unknown	<i>CASQ2</i>	NM_001232.3	c.872_880delTTGCCCGGG	p.Val291_Arg294del	deletion	homozygous	novel variant	N/A	N/A	P
46	M	23	Ph(+)-SCA	HCM	exercise-related	<i>TNNI2</i>	NM_001001430.2	c.275G>T	p.Arg92Gln	missense	heterozygous	novel variant	N/A	N/A	P
47	M	35	Ph(+)-SCA	ARVC	unknown	<i>PKP2</i>	NM_001005242	c.148_151delACAG	p.Thr50Serfs*61	frameshift	heterozygous	rs397516997	N/A	N/A	P
48	M	14	Ph(+)-SCA	DCM	unknown	<i>LMNA</i>	NM_170707.3	c.1621C>T	p.Arg541Cys	missense	heterozygous	rs56984562	N/A	N/A	P
49	M	14	Ph(+)-SCA	HCM	exercise-related	<i>TNNI3</i>	NM_000363	c.485G>C	p.Arg162Pro	missense	heterozygous	rs397516354	0.00005302	N/A	P
50	M	31	Ph(+)-SCA	DCM	exercise-related	<i>CAV3</i>	NM_001234.4	c.*6+2T>G	p.(?)	splice region	heterozygous	novel variant	N/A	N/A	LP
						<i>MYH6</i>	NM_002471.3	c.4193G>A	p.Arg1398Gln	missense	heterozygous	rs150815925	0.0005653	0.0004	VUS
51	M	63	Ph(+)-SCA	DCM	exercise-related	<i>DSP</i>	NM_004415.2	c.5326G>T	p.Glu1776*	stop-gain	heterozygous	novel variant	N/A	N/A	P
						<i>EYA4</i>	NM_172103	c.1340G>T	p.Gly447Val	missense	heterozygous	novel variant	N/A	N/A	VUS
52	M	31	Ph(+)-SCA	DCM	exercise-related	<i>DSP</i>	NM_001008844	c.3562T>C	p.Tyr1188His	missense	heterozygous	rs141508330	0.0001401	0.000067	LP
						<i>FLNC</i>	NM_001127487	c.5199+1G>T	p.?	splice region	heterozygous	rs1465588989	0.00006484	N/A	P
						<i>JPH2</i>	NM_020433	c.1210G>C	p.Ala404Pro	missense	heterozygous	novel variant	N/A	N/A	LP

53	M	43	Ph(+)/SCA	DCM	unknown	<i>TNXB</i>	NM_019105.6	c.10990G>A	p.Val3664Met	missense	heterozygous	novel variant	N/A	N/A	LP
54	M	9	Ph(+)/SCA	possible ARVC	unknown	<i>TTN</i>	NM_001256850	c.44440dupA	p.Thr14814Ansf7	frameshift	heterozygous	novel variant	N/A	N/A	LP
						<i>DSG2</i>	NM_001943.2	c.877A>G	p.Ile293Val	missense	heterozygous	rs2230234	0.08733	0.0687	VUS
						<i>DSP</i>	NM_004415.2	c.6208G>A	p.Asp2070Asn	missense	heterozygous	rs41302885	0.005636	0.0039	VUS
55	M	42	Ph(+)/SCA	MVP	exercise unrelated	<i>TNXB</i>	NM_019105.6	c.8734G>A	p.Gly2912Ser	missense	heterozygous	novel variant	N/A	0.0007	VUS
						<i>MYLK</i>	NM_053025.3	c.4169A>C	p.Asn1390Thr	missense	heterozygous	novel variant	N/A	N/A	VUS
						<i>HCN4</i>	NM_005477.2	c.2800C>T	p.Arg934Cys	missense	heterozygous	rs199638465	0.00008081	0.0006	VUS
56	M	40	Ph(+)/SCA	HCM	unknown	<i>TMPO</i>	NM_003276	c.689G>A	p.Arg230His	missense	heterozygous	rs758666714	0.00002324	0.000025	VUS
						<i>AKAP9</i>	NM_005751	c.9830T>C	p.Ile3277Thr	missense	heterozygous	rs144021475	0.0001937	0.000008	VUS
						<i>GJD4</i>	NM_153368	c.259C>T	p.Leu87Phe	missense	heterozygous	rs76906304	0.01812	0.013	VUS
57	F	17	Ph(+)/SCA	suspected SQTs	unknown	<i>CACNA2D1</i>	NM_000722	c.2264G>C	p.Ser755Thr	missense	heterozygous	rs151327713	0.0006633	0.0008	VUS
58	M	47	Ph(+)/SCA	MVP	exercise unrelated	<i>MYH6</i>	NM_002471	c.2383C>T	p.Arg795Trp	missense	heterozygous	rs202120238	0.0002014	0.000074	VUS
59	M	50	Ph(+)/SCA	possible ARVC	exercise-related	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
60	M	59	Ph(+)/SCA	DCM	exercise unrelated	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A

Abbreviations: ACMG, American College of Medical Genetics and Genomics; ARVC, arrhythmogenic right ventricular cardiomyopathy; DCM, dilated cardiomyopathy; HCM, hypertrophic cardiomyopathy; LP, likely pathogenic; LVNC, left ventricular non-compaction; MVP, mitral valve prolapse; P, pathogenic; Ph, phenotype; SCA, sudden cardiac arrest; SD, standard deviation; SQTs, short QT syndrome; VUS, variant of uncertain significance. N/A denotes 'not applicable'.

* *FKTN*-mediated diseases (DCM or muscular dystrophy-dystroglycanopathy) are transmitted as an autosomal-recessive trait; therefore, the identified heterozygous pathogenic variant is not considered to have a direct link to the SCA.

** Indicates patients who were successfully resuscitated, but died during hospitalization due to neurological complications caused by prolonged cerebral ischemia

† This variant has been initially interpreted as pathogenic; however, the recent evidence suggests contradictory results, and the variant may also be considered as VUS.

‡ This variant has been initially interpreted as pathogenic based on the available literature, but has been later reclassified as a VUS after additional *in vitro* patch clamp experiments and segregation studies revealed no evidence of pathogenicity.²⁶